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Effect of 4 weeks daily wild blueberry supplementation on symptoms of depression in adolescents

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Abstract:

Adolescence is an important period for cognitive maturation and emotional regulation and this age group is particularly vulnerable to developing depression. Diets rich in fruits and vegetables have been associated with decreased risk of developing depressive disorders across the lifespan, an association that may be due to the high flavonoid content of these foods. Previously we have shown increases in transient positive affect in both children and young adults two hours after administration of a wild blueberry intervention. Here, using a randomized double-blind, placebo-controlled trial, we investigated the effects of four weeks, daily wild blueberry supplementation (containing ~253mg anthocyanins) on transient and chronic mood in adolescents. Healthy 12-17-year old (N = 64, 35 females) were recruited and randomly assigned to receive either a wild blueberry or matched placebo supplementation. Depression and anxiety symptoms were assessed before and after the intervention period using the Mood and Feelings Questionnaire and Revised Child Anxiety and Depression Scale. Transient affect was assessed before, two weeks, and at four weeks using the Positive and Negative Affect Schedule. Following the intervention period there were significantly fewer self-reported depression symptoms in participants who were supplemented with the wild blueberry intervention compared to those who received the matched placebo ($p=0.02$, 95% CI -6.71 to -5.35). There was no between group effect on anxiety symptoms or on transient affect. Further investigation is required to identify specific mechanisms that link flavonoids consumption and mood. If replicated, the observed effects of wild blueberry supplementation may be a potential prevention strategy for adolescent depression and may have benefits for public mental health.

Introduction:

Puberty is a complex biologically driven process that has an impact on emotional and behavioral wellbeing, resulting in a period with increased risk of developing emotional disorders and risk-taking behavior. The brain undergoes cognitive maturation via synaptic remodeling well into the 20s. The limbic system, responsible for governing reward processing, appetite and pleasure seeking, matures before the prefrontal cortex, which is responsible for executive functioning such as problem solving, planning, emotional regulation and multitasking. This difference in cortical maturity is hypothesized to create a developmental imbalance, making teens vulnerable to behavioral and mental health problems, such as depression ⁽¹⁾.

An episode of major depressive disorder (MDD) during adolescence is a major personal and public health problem across the world ⁽²⁾. The disorder has many acute and long-term adverse consequences on adolescents' education and occupational success, relationships and family life and on their future physical and mental health ⁽³⁾. Each year around 7.5% of adolescents aged 13 to 18 years' experience an episode of MDD ⁽⁴⁻⁶⁾. Symptoms of MDD are distressing and include sleep and cognitive problems, low mood, irritability, feelings of worthlessness and lack of pleasure ⁽⁷⁾. Sub-clinical MDD is even more common: recent surveys in the UK suggest that ~25% of young people report elevated symptoms of depression in any given year ^(4,8), including depressive symptoms that are not sufficient in number or severe enough to meet diagnostic criteria. Sub-clinical symptoms have a major impact on daily functioning and are associated with increased risk of developing the disorder ⁽⁴⁾.

Treatment for MDD in this age group includes psychological therapies and anti-depressant medication; however, these are only moderately effective and are often inaccessible to young people due to limited public health service resources ⁽⁸⁾. A recent meta-analysis of psychological treatments for children and young people with mental health problems found that the effect size of treatment for depression was small ($d = 0.29$) and was lower than effects of treatment for other common mental health problems ⁽⁹⁾. Many young people with MDD do not receive an evidence-based treatment and the prevention of adolescent depression is, therefore, a highly valued goal ⁽¹⁰⁾. One potential way to prevent the onset of MDD and sub-clinical depression is through diet. Diet and depression symptoms are significantly associated in adults, although this relationship is complex and potentially bidirectional, i.e. unhealthy diet leading to low mood and vice versa ⁽¹¹⁾. A recent systematic review of the association between depression symptoms and diet in adolescents found that 'healthy' diets (i.e. consumption of fruits and vegetables) were associated with lower

79 depression symptoms; whilst ‘unhealthy’ diets (i.e. consumption of junk foods and saturated fats)
 80 were associated with higher depression symptoms ⁽¹²⁾. A large well-controlled epidemiological
 81 study examining associations between habitual intakes of dietary flavonoids and depression risk
 82 showed that individuals consuming diets higher in flavonoids presented a lower depression risk,
 83 particularly amongst older women ⁽¹³⁾. A similar study assessed symptoms of depression and the
 84 total habitual intake of polyphenols among the participants and found that higher dietary intake of
 85 flavonoids was inversely associated with depressive symptoms ⁽¹⁴⁾. Thus, diets rich in fruits and
 86 vegetables are associated with low depression symptoms. Dietary flavonoids are present in
 87 substantial concentrations in commonly consumed fruits and vegetables and may be a potential
 88 mediator for the anti-depressant action of diets rich in fruits and vegetables.

89
 90 The hypothesis that there is a causal relationship between diet and depression symptoms and the
 91 onset of MDD has recently been strengthened by number of intervention studies. Acute purple
 92 grape juice intervention resulted in increase in self-reported ratings of ‘calm’ in healthy young
 93 adults ⁽¹⁵⁾. Similarly, acute consumption of flavonoid-rich wild blueberry improved short-term
 94 positive mood in children aged 7-10 years and in young adults aged 18-25 years ⁽¹⁶⁾. In a recent
 95 randomized controlled trial with 67 depressed adults ⁽¹⁷⁾, participants randomized to an intervention
 96 promoting a healthy diet with at least nine portions of fruit and vegetables each day reported
 97 significantly less depression symptoms at twelve weeks than those randomized to receive social
 98 support. Anti-depressive effects of flavonoid rich plants and their extracts have also been
 99 investigated. *Hypericum perforatum* (also known as Saint John’s wort, derived from a flowering
 100 plant in the Hypericaceae family) extract intervention studies show its effectiveness as treatment for
 101 mild/moderate depression when compared to placebo and have similar effects to pharmacological
 102 treatments ⁽¹⁸⁻²¹⁾. Similarly, saffron (*Crocus sativus*, derived from the saffron spice of the flowering
 103 plant of *Crocus* genus) extract consumption had equivalent effect as pharmacological treatment for
 104 depression and was significantly more effective than the matched placebo ⁽²²⁻²⁴⁾.

105
 106 The specific effects of sustained wild blueberry flavonoid consumption on symptoms of depression
 107 in adolescents have not yet been tested. Here, we designed a double-blind, placebo-controlled
 108 experiment to test the effect of consuming a flavonoid-rich wild blueberry intervention for four
 109 weeks on symptoms of depression, anxiety and transient affect in healthy adolescents. Participants
 110 were randomly assigned to a wild blueberry or a matched placebo drink with transient affect and
 111 symptoms of depression and anxiety assessed before and after the four-week intervention period.

112

113

114 **Method**115 *Ethics*

116 This research was reviewed and given a favorable ethical opinion for conduct by the University of
117 Reading Research Ethics Committee (UREC 16/55). The study was registered at clinicaltrials.gov
118 NCT03119597.

119

120 *Participants*

121 An *a priori* power analysis (using G Power 3.1.9.2) based on data from a previous study⁽¹⁶⁾
122 revealed that 24 participants per group were required to achieve power of 0.8 with alpha set at 0.5
123 level. Students aged 11-17 years of varying ethnicity, from four schools in Reading Berkshire, UK
124 were invited to take part in this study. All parents or legal guardians provided informed written
125 consent for young people under the age of 16. Participants under the age of 16 provided written
126 assent and those over 16 gave written consent. All participants were screened for any health
127 conditions (including mental health), any treatment they were receiving and food related allergies
128 that would exclude them from the study. We screened 82 young people, of whom 18 dropped out
129 after the first screening session. Sixty four participants were randomly assigned to either a wild
130 blueberry drink or a matched placebo drink. The randomized allocation of participants to treatment
131 was generated using excel. The groups were coded A and B and the sequence was saved in a
132 password protected spreadsheet. Both the researchers and the participant were blind to treatment
133 group and participants were told the study was investigating effects of different fruit drinks so were
134 not aware of the study hypothesis.

135

136 *Interventions*

137 Both interventions (wild blueberry and placebo) were measured and packaged into silver opaque
138 sachets at the University of Reading. Sachets were identical for the wild blueberry and the placebo
139 drink and neither the researchers nor the participants knew what their sachets contained. Wild
140 Blueberry Association of North America provided the blueberry powder whilst the matched sugars
141 and vitamin C (placebo) was obtained from Bulk Powders. The packets of wild blueberry contained
142 13g of freeze-dried wild blueberry (WBB) powder (containing ~253mg anthocyanins). Placebo
143 packets were matched to the WBB for sugars (4.52g glucose and 4.79g fructose) and vitamin C (4
144 mg). Each participant was given 14 days' supply of their requisite intervention, along with written
145 and video instructions for their parents/guardians on how to prepare the intervention. Each
146 intervention was prepared daily, by adding 30 ml of low-flavonoid 'Rock's Organic Orange

Squash' and 170 ml of water and the contents of the sachet to the opaque cup provided. Each participant was given a checklist to record the dates and times when they consumed the drink each day and the name of the person who prepared the drinks. Participants were also asked to bring back their used sachets after two weeks as a measure of compliance. The remaining 14 days' supply of each intervention was given to the participants two weeks into the intervention period. The true aim of the study was not disclosed to the participants, they were informed that it was a fruit drink study, to avoid revealing the contents of the drink.

Measures

The Mood and Feelings Questionnaire (MFQ) was used to measure symptoms of depression ⁽²⁵⁾. The MFQ is considered to be the gold standard self-report measure for depression in young people (NICE, 2015). It is a standardized and well-validated 33-item self-report measure of the severity of depression symptoms in adolescents. Each item relates to a symptom or experience associated with depression. Participants are asked to rate each item in relation to their symptoms in the past 2 weeks on a 3-point Likert scale (not true = 0, sometimes = 1, true = 2). Total MFQ scores range from 0 to 66 where higher scores indicate greater risk of depression. The clinical cut off for the MFQ is 27, with scores above 27 indicating significant risk of a diagnosis of MDD ⁽²⁵⁾.

Anxiety symptoms were assessed using the anxiety sub-scale of the Revised Child Anxiety and Depression Scale (RCADS) ⁽²⁶⁾, a standardized and validated measure of anxiety symptoms in young people used routinely in UK NHS mental health services. The anxiety sub-scale of RCADS consists of 37 items, each rated on a 4-point Likert scale (never = 1, sometimes = 2, often = 3, always = 4). Total scores range from 37 to 148 with higher scores indicating increased risk of an anxiety disorder. Again, participants were asked to rate the items keeping the past two weeks in mind.

Current mood (i.e. transient affect) was assessed using the Positive and Negative Affect Schedule-NOW (PANAS-NOW) at screening, and at two and four weeks. As the term suggests this is a measure of transient mood. The PANAS is a valid and reliable 20 self-report measure of positive affect (PA – 10 items) and negative affect (NA - 10 items) that can be used on multiple test occasions ^(27,28). Participants rated the degree to which they were currently experiencing each item on a 5-point Likert scale ranging from 'very slightly' to 'extremely'. Ratings of positive and negative items were summed to calculate an overall positive affect and overall negative affect score, each ranging from 10-50 where lower scores indicate lower levels of positive or negative affect.

181

182 Habitual fruit and vegetable consumption were assessed using EPIC-Norfolk food frequency
 183 questionnaire, a semi-quantitative paper-based questionnaire, which includes 130 food items, each
 184 rated on 9-point Likert scale (never or less than a month-1 to 6+perday-9). FETA software was used
 185 to analyse the data collected to calculate 46 nutrient and 14 food group values including average
 186 daily fruit and vegetable intake ⁽²⁹⁾.

187

188 *Other measures i.e. working memory, verbal fluency, cognitive accuracy and reaction time were*
 189 *assessed and are reported elsewhere* ⁽³⁰⁾.

190

191

192 *Procedure*

193 As outlined in Figure 1, participants were seen by the researchers four times across a five weeks
 194 period. All participants did not attend all assessment – the number of participants assessed at each
 195 timepoint is indicated in Figure 1. Research sessions took place either at the University of Reading
 196 or at the participant's school. Sessions were scheduled at the same time of day for each participant.
 197 The first two sessions, scheduled 48 hours apart, were screening sessions where participants
 198 completed a battery of questionnaires: MFQ, RCADS, (screening session 1) PANAS, EPIC-
 199 Norfolk food frequency questionnaire and a questionnaire about their health status (screening
 200 session 2). Screening sessions were limited to 30 minutes to fit with the school timetable and to
 201 maintain high levels of participant engagement in both sessions. Parents were also asked to
 202 complete a demographic questionnaire. Participants started the intervention the day after the
 203 second screening session was completed. Two weeks later they returned their used drink sachets,
 204 were given a new checklist and completed the PANAS (Test session 1). Participants were also
 205 asked if they were experiencing any adverse effects of the drink and feedback on its palatability.
 206 They then returned two weeks later (Test session 2), returned their drink sachets, completed the
 207 PANAS, MFQ and RCADS and were debriefed. For each test session, participants were instructed
 208 not to consume their allocated intervention before the test session to ensure that chronic, not acute,
 209 effects of the intervention were being measured.

210

211 *Statistical Analysis*

212 Statistical analyses were conducted using IBM SPSS version 22. T-test was used to investigate
 213 differences in symptoms of depression, anxiety and fruit and vegetable intake between the two
 214 groups at baseline. Effects of intervention on transient affect was analysed using Linear Mixed

Modelling (LMM) using an unstructured covariance matrix to model successive repeat test sessions, with subjects included as random effects. Data from two weeks and four weeks measures of the PANAS and treatment group were included as fixed factors, with baseline PANAS scores included as a covariate. LMM deals with data that is missing at random and with multiple measurement points, giving unbiased estimates of each of the means. To test the effects of the intervention on anxiety and depressive symptoms at four weeks, data were analysed using Analysis of Covariance (ANCOVA) with drink (Placebo, WBB) as an independent variable and MFQ and RCADS scores at 4 weeks as dependent variable. Baseline measures of depression and anxiety were used as covariates and Bonferroni corrected t-tests were used to investigate all fixed effects and interactions.

Results

Sample characteristics

Sixty-four participants were randomised (35 females, 29 males) aged 12-17 years ($M = 14.20$ years, $SD = 1.71$). Thirty-five participants were randomly allocated to receive the placebo drink and twenty-nine to the WBB intervention. Participants' demographic data, baseline mood scores and habitual fruit and vegetable intakes are reported in table 1. There were no significant differences between groups in the amount of daily fruit $t(51) = 0.14$, $p = 0.89$ or vegetables $t(51) = 1.45$, $p = 0.15$ consumed. One sample t-test revealed that the mean fruit and vegetable consumption by the participants was significantly lower than the 400g per day as recommended by WHO; fruit: $t(52) = 11.20$, $p < 0.005$, vegetables $t(52) = 7.12$ $p < 0.005$.

At baseline mean depression and anxiety scores were 12.35 ($SD = 9.31$) and 23.19 ($SD = 13.80$) respectively, both below the clinical threshold. There was no significant group difference in symptoms at baseline; MFQ $t(60) = 0.60$, $p = 0.55$, RCADS $t(40) = 0.45$, $p = 0.66$ and no group difference in mean positive and negative affect; $t(62) = 1.40$, $p = 0.17$ and $t(62) = 0.80$, $p = 0.98$ respectively. A minority of participants (9.38%) reported depression symptoms above the clinical cut-off of 27 on the MFQ (11.4% in the placebo group, 3.4% in the intervention group). No participants reported anxiety symptoms above the clinical threshold. No participants reported a diagnosis of depression or anxiety, or that they were receiving treatment for these disorders.

Hypothesis testing

At four weeks 59 participants provided self-report data on anxiety (RCADS) and depression (MFQ) symptoms; 26 from the intervention group and 33 from the placebo group. As shown in Figure 2a,

after four weeks of the intervention, the mean MFQ score for participants who consumed WBB was significantly lower than the mean MFQ score for participants who consumed the placebo drink. This was significant $F(1,57)=5.52$, $p=0.02$ 95%CI -6.71 to -5.35 with a medium effect size ($d = 0.65$). The change in the depression scores for each participant including regression line for both treatments is shown in figure 3. There was no significant effect of WBB on symptoms of anxiety (Figure 2b) after four weeks of supplementation $F(1,34) = 2.1$, $p=0.16$; mean RCADS score for participants in the WBB group was 13.90, (SD = 8.39) and the mean RCADS for the placebo group was 19.3, (SD = 11.31).

We also examined the effect of intervention on positive affect and negative affect (PANAS) after two and four weeks (see Figure 4). There was no significant effect of Drink, $F(1,64.33) = 0.26$, $p=0.62$, Repeated trial, $F(1,62.22) = 2.95$, $p=0.09$, or any Drink x Repeated trial interaction $F(1,62.22) = 3.686$, $p=0.06$ on transient positive affect. Figure 4a shows the mean PA scores following intervention of WBB and placebo at two weeks and at four weeks. There was also no significant effect of the intervention on NA; Repeated trial, $F(1,59.3) = 0.66$ $p=0.42$, Drink, $F(1,63.79) = 0.24$ $p=0.63$ or Repeated trial \times Drink interaction, $F(1,59.30) = 1.17$, $p=0.28$. As shown in Figure 4b, NA was not significantly different after consuming the WBB drink or the placebo drink.

Discussion

This randomized, placebo controlled, double blinded trial investigated the effects of 4 weeks consumption of a flavonoid-rich WBB drink on symptoms of depression and anxiety and on transient affect in a community sample of healthy 12-17-year old. The results demonstrated that after four weeks of daily WBB intervention there was a between groups difference in self-reported depressive symptoms; participants randomised to the WBB intervention reported significantly lower scores on the measure of depression symptoms than participants who were randomised to the placebo drink. There was no significant effect of the intervention on anxiety symptoms or on positive affect or negative affect (i.e. transient affect). The data suggest that flavonoid supplementation may be beneficial in reducing depressive symptoms in healthy adolescents.

This is, to our knowledge, the first randomized double blinded study to show the effects of chronic WBB flavonoids on depression symptoms in teenagers. The participants in the study were healthy but at baseline assessment were consuming sub-optimal habitual levels of flavonoids, i.e. their daily consumption of fruit (44.87%) and vegetable (57.46%) was well below the WHO recommended

283 amount of 400g/day ^(31,32). This is consistent with the typical diet of young people in the UK, where
 284 only 18% of adolescents meet the recommended daily requirement, and the average daily
 285 consumption within this age group is 256g (3.5 portions) of fruit and vegetables ⁽³³⁾. Levels of
 286 depression and anxiety were similar to community norms on gold standard self-report measures.
 287 Importantly, because the effects of the intervention were observed in a community sample, these
 288 effects cannot necessarily be generalised to adolescents with more severe symptoms of depression
 289 or a diagnosis of depression.

290

291 Within this community sample the effect size of the flavonoid intervention compared to the control
 292 group on the measure of depression symptoms, the MFQ, was $d = 0.65$, a medium effect size. To
 293 put this into context, two recent meta-analyses have examined the effects of psychological
 294 treatments for depression and the prevention of depression. Ecksthtain et al., (2019) concluded that
 295 the treatment effect size of psychological treatments for adolescents with depression was $d = .36$
 296 ⁽³⁴⁾. In a review of interventions to prevent depression Ssegonia et al., (2019) reported an effect size
 297 of $d = .22$ ⁽³⁵⁾. In relation to the specific measure of depression used in this study the reduction of
 298 the 4 points on mean MFQ scores in the intervention group indicates complete amelioration of 2
 299 items on the scale or a reduction (from 2 to 1, or 1 to 0) of 4 items. Because each item reflects a
 300 symptom or adverse effect of depression, clinically this would be likely to reflect a meaningful
 301 reduction in the impact of depression on the young person ⁽³⁶⁾.

302

303 Previously the effects of flavonoids from different sources such as apples, cocoa and grape juice
 304 showed no effects on depression in healthy adults ⁽³⁷⁻⁴⁰⁾. However, our results are consistent with
 305 previous animal and epidemiological studies that suggest anti-depressive effects of a flavonoid rich
 306 diet ^(13,41-44). They also are in keeping with experimental data on the acute effects of WBB on
 307 positive mood in children and young adults ^(15,16), and the acute effect of grape juice on mood in
 308 healthy adults ⁽⁴⁵⁾. Unlike a previous acute intervention study, we did not observe a significant
 309 effect of WBB on momentary mood (i.e. transitory affect). However, the interval between
 310 consuming the WBB drink and assessing NA and PA was variable, unlike the standard 2-hour
 311 interval used in previous studies. In addition, the four-week assessment (our end point) was
 312 conducted during the first week of school after the summer holidays. Unlike symptoms of
 313 depression (and anxiety) which were measured over a minimum two-week period and which are
 314 conceptualised as relatively stable, positive and negative affect are conceived as short-lived events
 315 that have rapid decay after elicitation ⁽⁴⁶⁾. It is therefore possible that this external event (returning
 316 to school) had a measurable impact on participants' momentary affect.

317

318 Although anxiety and depression are frequently co-morbid in young people and share some
 319 symptoms (e.g. fatigue, low concentration and sleep disturbances), the results of this intervention
 320 study suggest that flavonoids may reduce symptoms that are more prominent in depression than
 321 anxiety, e.g. low mood, anhedonia, feelings of guilt, and worthlessness and do not reduce symptoms
 322 that are specific to anxiety. However, it is also possible that the effect of flavonoids on anxiety is
 323 smaller than the effect on depression and that a larger sample, with greater power, might result in a
 324 significant effect.

325

326 Some authors have proposed that flavonoids increase cerebral blood flow to the dorsolateral
 327 prefrontal cortex, a site that is highly associated with cognitive and emotional regulation, including
 328 rumination, a cognitive process of repetitive thinking that may exacerbate feelings of guilt and
 329 worthlessness⁽⁴⁷⁻⁴⁹⁾. This suggests that there may be an indirect pathway between flavonoid
 330 consumption and depression whereby flavonoid consumption enhance cerebral blood flow, which
 331 boosts executive functioning; in turn improved executive functioning helps to enhance cognitive
 332 control, inhibits rumination and thus reduces depression. Adolescents with depression have
 333 impaired executive function compared to non-depressed and anxious young people⁽⁵⁰⁾ and therefore
 334 the benefits of flavonoid consumption may be more prominent in these young people. However,
 335 potentially any positive effects of flavonoid consumption on executive function would have benefits
 336 for more young people because executive function is critical for academic achievement⁽⁵¹⁾.

337

338 A plausible direct pathway between flavonoid consumption and mood is the effects of flavonoids
 339 on Monoamine Oxidase (MAO)⁽⁵²⁾. MAO inhibitors have been used to treat mood disorders and
 340 flavonoids may mimic their effects^(52,53). A recent study showed that consuming fruits high in
 341 flavonoids i.e. blackcurrants significantly reduces MAO activity and increases the circulating
 342 monoamines and thereby elevates mood⁽⁵²⁾. Another possible mechanism by which flavonoids may
 343 affect mood is by mimicking anxiolytic-like effects by binding to benzodiazepine receptors,
 344 enhancing the effect of GABA via GABAA receptors^(34,54,55). However, in line with a previous
 345 study⁽¹⁶⁾ that showed no changes in negative affect (an indicator of anxiety) after acute flavonoid
 346 intervention, here there was no significant of flavonoid consumption on anxiety.

347

348 Although the mechanisms of action require further investigation there is accumulating evidence of a
 349 causal relationship between flavonoid consumption and depression symptoms. This evidence has
 350 been published by independent research groups using different research designs, including

epidemiology, clinical trials and experiments. However, the research is preliminary and requires robust replication and extension, with larger samples, longer time scales and careful tests of mechanisms of action. Our study examined the effects of flavonoids on healthy young people, some of whom had elevated symptoms of depression. We did not have adequate power to conduct subgroup analysis but clearly it is important to identify if the change in depression symptoms is driven by improvements in those with relatively elevated symptoms, or if the effects are similar across all levels of baseline depression. This distinction is important because flavonoids may have the potential to prevent depression in those at risk (i.e. those with elevated symptoms) or may have a more general effect. The former would suggest that dietary interventions could be used for early intervention in those exhibiting symptoms of depression; the latter that dietary interventions could have a broader benefit to public mental health.

Conclusions

This randomized double-blind study demonstrated the chronic effects of wild blueberry flavonoid consumption on reducing symptoms of depression in a community sample of adolescents. Dietary flavonoid interventions may have potential to reduce symptoms of depression in adolescents. This study requires replication, not only in healthy participants, but also in clinically referred samples to assess the potential of flavonoids to be used as a practical and cost-effective intervention. In addition to this, studies focused on investigating biochemical changes and investigating the mechanistic pathways in which flavonoids decrease depressive symptoms in humans is essential.

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Authorship

All the authors were involved in the design of the experiments; S.K, and J.F performed the experiments and analysed the data. S.K, J.F, C.W and S.R were involved in the writing and revisions of the manuscript.

385 **Conflicts of Interest**

386 The authors declare no conflicts of interest arising from the conclusions of this work.

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534 **TABLES**

535 Table 1: Demographic details, mean fruit and vegetable intake and mean depression and anxiety
 536 scores at baseline for both intervention groups.

	PLACEBO GROUP	WILD BLUEBERRY GROUP	P VALUES
MEAN AGE	14.5 (SD=1.804)	13.82(SD=1.54)	P=0.11
MALE %	48.6	41.4	P=0.57
FEMALE %	51.4	58.6	P=0.57
BRITISH %	60	52.4	P=0.52
ASIAN%	11.4	12.5	P=0.52
MIXED%	5.8	12.6	P=0.52
AFRICAN	2.9	8.3	P=0.52
CHINESE	2.9	4.2	P=0.52
MEAN FRUIT INTAKE (GRAMS/DAY)	188 (SD=168.3)	176 (SD=98.0)	P=0.89
MEAN VEGETABLES (GRAMS/DAY)	257.6 (SD= 187.0)	187.5 (SD=144.6)	P=0.15
MEAN DEPRESSION (MFQ)	13.0 (SD= 10.0)	11.3 (SD= 8.5)	P=0.55
MEAN ANXIETY (RCADS)	24.2 (SD= 14.90)	22.3 (SD=13.0)	P=0.66
MEAN POSITIVE AFFECT	28.0 (SD=7.7)	25.3 (SD=8.0)	P=0.17
MEAN NEGATIVE AFFECT	15.1 (SD=5.24)	14.1 (SD=4.38)	P=0.98

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540 **FIGURES**

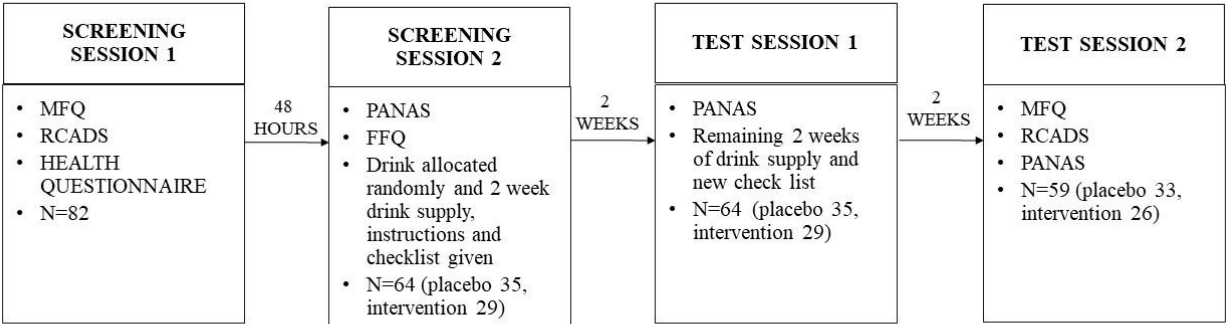


Figure 1. A schematic of the study procedure

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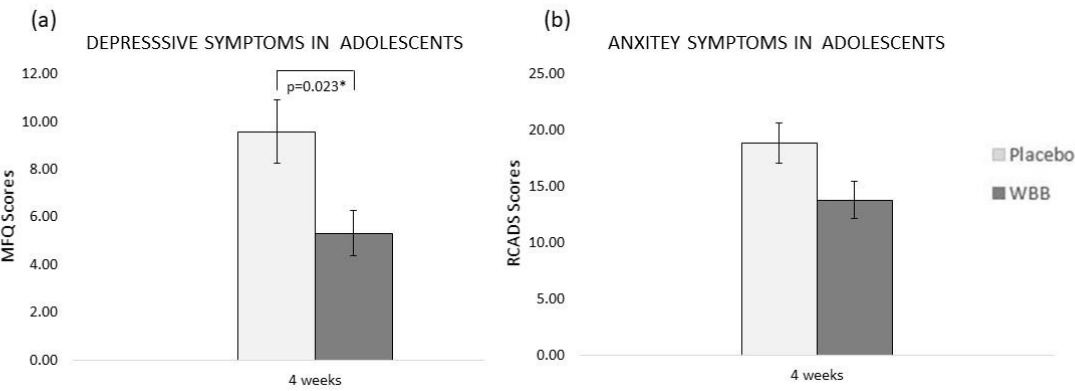


Figure 2. Mean scores (\pm standard error of the mean) in adolescents aged 11-17 years (a) Mean MFQ scores after 4 weeks consumption of placebo and intervention drinks. (b) Mean RCADS scores after 4 weeks consumption of placebo and intervention drinks.

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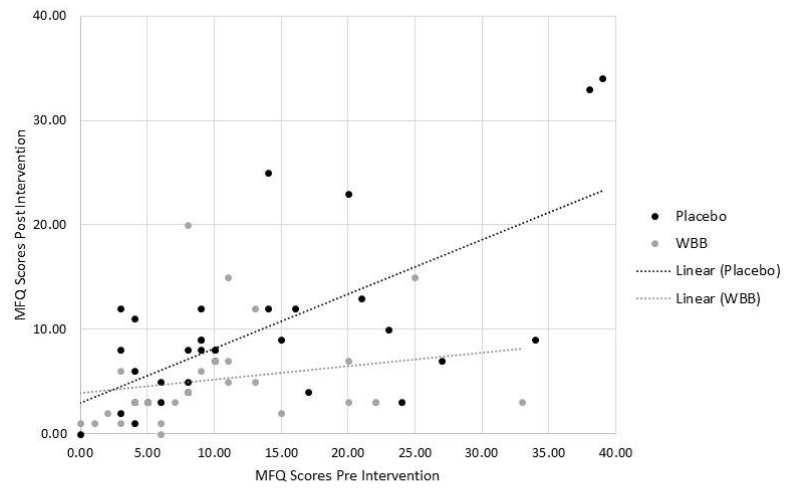


Figure 3. Scatterplot showing the MFQ scores at baseline and 4-week post intervention

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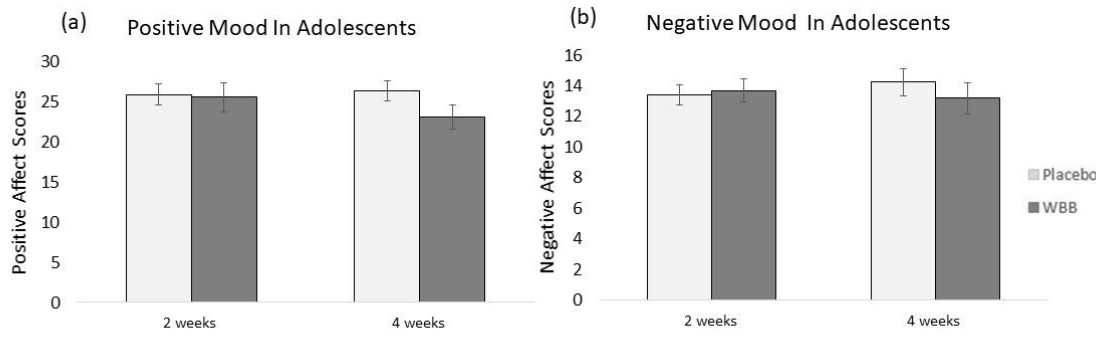


Figure 4. Mean PANAS-NOW Mood scores (\pm standard error of the mean) in adolescents aged 11-17 years: (a) Mean PA scores 2 and 4 weeks post-consumption of placebo and intervention drinks. (b) Mean NA scores 2 and 4 weeks post-consumption of placebo and intervention drinks.

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